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  => s 12 (P) clot or thromb?
L3 652552 L2 (P) CLOT OR THROMB?
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         Dot Format:
                                            DD.MM.YYYY or MM.YYYY
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Feb. 10, 1987
Feb. 10, 2000
Feb 10, 1987
         Text Format:
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March 5 - 8, 1990
April - June, 1999
                                            10 February 1987
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        Any year entered with only two digits will be interpreted as being in the range 1900-1999. Thus, Mar 12 01 will be searched as
        19010312.
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3 FILES SEARCHED...
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L12 ANSWER 1 OF 7 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1999:794322 CAPLUS
 DOCUMENT NUMBER:
                                                            132:18789
 TITLE:
                                                            Compositions and methods using an oxidized/reduced
                                                            low-molecular-weight heparin compound for inhibiting
                                                             thrombogenesis
                                                            Hirsh, Jack; Weitz, Jeffrey I.
 PATENT ASSIGNEE (S):
                                                            Hamilton Civic Hospitals Research Development Inc.,
                                                            Can.
SOURCE:
                                                           U.S., 48 pp., Cont.-in-part of U.S. 5,763,427. CODEN: USXXAM
 DOCUMENT TYPE:
                                                            Patent
LANGUAGE:
                                                            English
 FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
            PATENT NO.
                                                    KIND DATE
                                                                                                       APPLICATION NO.
            US 6001820
                                                                  19991214
                                                                                                      US 1997-870528
                                                                                                                                              19970606
                                                                                                     US 1997-870528
US 1995-540324
AU 1996-51400
US 1996-624327
JP 1996-528734
NO 1997-4500
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                                                                                                                                              19951006
19960329
                                                                                                             1996-624327
         WO 1996-CA190 19960329

R SOURCE(S): MARPAT 132:18789

Compns. and methods are provided for the treatment of cardiovascular diseases. More particularly, the invention relates to modifying thrombus formation by administering an agent which, inter alia, is capable of (1) selectively inactivating thrombin which is bound either to fibrin in a clot or to some other surface, but which has only minimal inhibitory activity against free thrombin, i.e., fluid-phase thrombin; (2) inhibiting the assembly of the intrinsic tenase complex, thereby inhibiting the activation of Factor IXa; and (3) inhibiting the activation of Factor IXa; and (3) inhibiting the activation of Factor IX by Factor XIa. The compns. and methods of the present invention are particularly useful for preventing thrombosis in the circuit of cardiac bypass app. and in
                                                                                                      WO 1996-CA190
                                                                                                                                              19960329
OTHER SOURCE(S):
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patients undergoing renal dialysis, and for treating patients suffering from or at risk of suffering from thrombus-related cardiovascular conditions, such as unstable angina, acute myocardial infarction (heart attack), cerebrovascular accidents (stroke), pulmonary embolism, deep vein thrombosis, arterial thrombosis, etc. The invention uses a polyanionic carbohydrate, esp. an oxidized/reduced low-mol.-wt. heparin compd. (prepn. described) REFERENCE COUNT: 57
(1) Alhenc-Gelas; Fundamental and Clinical Cardiology 1994, V19, P43 CAPLUS
(2) Anon; WO 8201005 1982 CAPLUS
(3) Anon; WO 8203627 1982 CAPLUS
(4) Anon; EP 101141 1984 CAPLUS
(5) Anon; EP 121067 1987 CAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
HD DATE APPLICATION NO. DATE REFERENCE (S): PATENT NO. KIND DATE APPLICATION NO. US 6001820 19991214 US 1997-870528 US 1995-540324 AU 1996-51400 19970606 A A1 US 5744457 19980428 19951006 <--AU 9651400 19961016 19960329 <--19960329 <--US 5763427 19980609 US 1996-624327 JP 1996-528734 JP 11506420 T2 19990608 JP 1996-524374 19960329 <-NO 9704500 T2 19990608 JP 1996-528734 19960329 <-NO 9704500 A 19971128 NO 1997-4500 19970929 <-Compns. and methods are provided for the treatment of cardiovascular
diseases. More particularly, the invention relates to modifying thrombus
formation by administering an agent which, inter alia, is
capable of (1) selectively inactivating thrombin which is bound either to
fibrin in a clot or to some other surface, but which has only minimal
inhibitory activity against free thrombin, i.e.,
fluid-phase thrombin; (2) inhibiting the assembly of
the intrinsic tenase complex, thereby inhibiting the activation
of Factor X by Factor IXa; and (3) inhibiting
the activation of Factor IX by Factor XIa. The
compns. and methods of the present invention are particularly useful for
preventing thrombosis in the circuit of cardiac bypass app. and in
patients undergoing renal dialysis, and for treating patients suffering
from or at risk of suffering from thrombus-related cardiovascular
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attack), cerebrovascular accidents (stroke), pulmonary embolism, deep vein
thrombosis, arterial thrombosis, etc. The invention uses a polyanionic
carbohydrate, esp. an oxidized/reduced low-mol.-wt. heparin compd. (prepn.
described). JP 11506420 19990608 19960329 <--37203-61-5, Blood coagulation factor XIa 37316-87-3, Blood coagulation factor IXa BAC (Biological activity or effector, except adverse); BIOL (Biological study)
(oxidized/reduced low-mol.-wt. heparin compd. for inhibiting thrombogenesis) L12 ANSWER 2 OF 7 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1999:619353 CAPLUS 1999:619353 CAPLUS 131:241741 DOCUMENT NUMBER: TITLE: A human antibody that binds to the .gamma.-carboxyglutamic acid domain of factor IX is a .gamma.-Carboxyglutamic acid domain of factor IX is potent antithrombotic in vivo Refino, Canio J.; Himber, Jacques; Burcklen, Louis; Moran, Paul; Peek, Mark; Suggett, Shelley; Devaux, Brigitte; Kirchhofer, Daniel Cardiovascular Research Antibody Technologies Dep., Genentech Inc., South San Francisco, CA, 94080, USA Thromb. Haemostasis (1991, 82(3), 1188-1195 CODEN: THHADO; ISSN: 0340-6245 AUTHOR(S): CORPORATE SOURCE: SOURCE: PUBLISHER: F. K. Schattauer Verlagsgesellschaft mbH JISHER: F. K. Schattauer Verlagsgesellschaft mbH MENT TYPE: Journal SUAGE: English

The human antibody F(ab')2, 10C12, which specifically binds to the GIa domain of factor IX, interfered with all known coagulation processes that involve factor IX/IXa. These include the function of the intrinsic Xase complex and the activation of zymogen factor IX by factor XIa and by the tissue factor:factor VIIa complex. 10C12 potently inhibited activated partial thromboplastin clotting times (APTT) in plasma of guinea pig and rat, thus enabling in vivo evaluation. In guinea pigs, a bolus administration of 10C12 (10 .mu.g/kg) prevented cyclic flow variations in damaged carotid arteries without affecting coagulation or bleeding parameters. At a 100-fold higher dose, 10C12 had no effect on normal hemostassis as assessed by the cuticle bleeding time. At this dose, 10C12 was also efficacious in a rat arterial thrombosis model, substantially reducing clot wt. and duration of vessel occlusion while prolonging ex vivo APTT only 1.2-fold. The dose of heparin required to produce comparable anti-thrombotic effects prolonged the APTT by 12-fold and increased the tail bleeding time (TBT) by 8-fold. In contrast, 10C12 had no effect on TBT. Rat tails showed a tendency for rebleeding which 10C12 exacerbated. In conclusion, the antithrombotic potency of the 10C12 exacerbated. In conclusion, the antithrombotic potency of the 10C12 antibody in 2 species provides evidence for an important role of FIX, and its GIa domain in particular, during thrombogenesis under arterial flow conditions. The relative safety at EDs of this fully human antibody suggests that it may have therapeutic value for treatment of thrombotic disorders. Journal LANGUAGE: REFERENCE COUNT: REFERENCE(S): (1) Ahmad, S; J Biol Chem 1989, V264, P20012 CAPLUS (2) Ahmad, S; Trends Cardiovasc Med 1994, V4, P271 CAPLUS (3) Baselga, J; J Clin Oncol 1996, V14, P737 CAPLUS
(4) Benedict, C; Blood 1993, V81, P2059 CAPLUS
(5) Benedict, C; J Clin Invest 1991, V88, P1760 CAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT Thromb. Haemostasis (1999), 82(3), 1188-1195 CODEN: THHADQ; ISSN: 0340-6245 CODEN: THHADO; ISSN: 0340-6245
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SO

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L12 ANSWER 3 OF 7 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1999:443190 CAPLUS
            ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                                                                                                                              131:208823
                                                                                                                                                             Targeted inhibition of intrinsic coagulation limits cerebral injury in stroke without increasing intracerebral hemorrhage
             TITLE:
          AUTHOR (S):
                                                                                                                                                             Choudhri, Tanvir F.; Hoh, Brian L.; Prestigiacomo,
Charles J.; Huang, Judy; Kim, Louis J.; Schmidt, Ann
Marie; Kisiel, Walter; Connolly, E. Sander, Jr.;
Pincky, David J.
                                                                                                                                                          Marie: Kisiel, Walter: Connolly, E. Sander, Jr.;
Pinsky, David J.
Department of Neurological Surgery, Columbia
University College of Physicians and Surgeons, New
York, NY, 10032, USA
J. Exp. Med. 1999, 190(1), 91-99
CODEN: JEMEAV, ISSN: 0022-1007
Rockefeller University Press
          CORPORATE SOURCE:
          SOURCE:
          PUBLISHER:
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English
          DOCUMENT TYPE:
                               MENT TYPE: Journal SUAGE: English

Agents that restore vascular patency in stroke area increase the risk of intracerebral hemorrhage (ICH). As Factor IXA is a key intermediary in the intrinsic pathway of toagulation, targeted inhibition of Factor IXA-dependent coagulation might inhibit microvascular thrombosis in stroke without impairing extrinsic hemostatic mechanisms that limit ICH. A competitive inhibitor of native Factor IXA for assembly into the intrinsic Factor IXA activation complex, Factor IXAi, was prepd. by covalent modification of the Factor IXA active site. In a modified cephalin clotting time assay, in vivo administration of Factor IXAi caused a dose-dependent increase in time to clot formation (3.6-fold increase at the 300 .mu.g/kg dose compared with vehicle-treated control animals, P < 0.05). Mice given Factor IXAi and subjected to middle cerebral artery occlusion and reperfusion demonstrated reduced microvascular fibrin accumulation by immunoblotting and immunostaining, reduced 111In-labeled platelet deposition (42% decrease, P < 0.05), increased cerebral perfusion (2.6-fold increase in ipsilateral blood flow by laser doppler, P < 0.05), and smaller cerebral infarcts than vehicle-treated controls (70% redn., P < 0.05) based on tri-Ph tetrazolium chloride staining of serial cerebral sections. At therapeutically EDs, Factor IXAi was not assocd. with increased ICH, as opposed to tissue plasminogen activator (tPA) or heparin, both of which significantly increased ICH. Factor IXAi was cerebroprotective even when given after the onset of stroke, indicating that microvascular thrombosis continues to evolve (and may be inhibited) even after primary occlusion of a major cerebrovascular tributary.
          LANGUAGE:
       tributary.
REFERENCE COUNT:
                                                                                                                                                        (1) Benedict, C; J Clin Invest 1991, V88, P1760 CAPLUS (4) Choudhri, T; J Clin Invest 1998, V102, P1301
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                                                                                                                                                                                CAPLUS
                                                                                                                                                    CAPLUS
(5) Choudhri, T; Stroke 1997, V28, P2296 CAPLUS
(6) Connolly, E; Circ Res 1997, V81, P304 CAPLUS
(7) Connolly, E; J Clin Invest 1996, V97, P209 CAPLUS
ALL CITATIONS AVAILABLE IN THE RE FORMAT
                                J. Exp. Med. (1999), 190(1), 91-99

CODEN: JEMANY; ISSN: 0022-1007

Agents that testope vascular patency in stroke also increase the risk of intracerebral nemorrhage (ICH). As Factor IXa is a key intermediary in the intrinsic pathway of coagulation, targeted inhibition of Factor IXa-dependent coagulation
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                           key intermediary in the intrinsic pathway of coagulation, targeted inhibition of Factor IXa-dependent coagulation might inhibit microvascular thrombosis in stroke without impairing extrinsic hemostatic mechanisms that limit ICH. A competitive inhibitor of native Factor IXa for assembly into the intrinsic Factor X activation complex, Factor IXa, was prepd. by covalent modification of the Factor IXa active site. In a modified cephalin clotting time assay, in vivo administration of Factor IXa; caused a dose-dependent increase in time to clot formation (3.6-fold increase at the 300 mu.g/kg dose compared with vehicle-treated control animals, P < 0.05). Mice given Factor IXa; and subjected to middle cerebral artery occlusion and reperfusion demonstrated reduced microvascular fibrin accumulation by immunoblotting and immunostaining, reduced 111In-labeled platelet deposition (42% decrease, P < 0.05), increased cerebral perfusion (2.6-fold increase in ipsilateral blood flow by laser doppler, P < 0.05), and smaller cerebral infarcts than vehicle-treated controls (70% redn., P < 0.05) based on tri-Ph tetrazolium chloride staining of serial cerebral sections. At therapeutically EDs, Factor IXai was not assocd. with increased ICH, as opposed to tissue plasminogen activator (tPA) or heparin, both of which significantly increased ICH. Factor IXai was cerebroprotective even when given after the onset of stroke, indicating that microvascular thrombosis continues to evolve (and may be inhibited) even after primary occlusion of a major cerebrovascular tributary.
L12 ANSWER 4 OF 7 CAPLUS COPYRIGHT 2001 ACACCESSION NUMBER: 1998:347694 CAPLUS DOCUMENT NUMBER: 129:117633
                                                                                                                                                              COPYRIGHT 2001 ACS
                                                                                                                                                   Heparinless cardiopulmonary bypass with active-site
                                                                                                                                              Heparinless cardiopulmonary bypass with active-site blocked factor IXA: a preliminary study on the dog Spanier, Talia B.; Oz, Mehmet C.; Minanov, Oktavijan P.; Simantov, Ronit; Kisiel, Walter; Stern, David M.; Rose, Eric A.; Schmidt, Ann Marie Department of Surgery, Calumbia_University College of Physicians and Surgeons, Naw York, NY, 10032, USA J. Thorac. Cardiovasc. Surg. (1998), 115(5), 1179-1188
 AUTHOR (S):
CORPORATE SOURCE:
SOURCE:
                                                                                                                                                CODEN: JTCSAQ; ISSN: 0022-5223
Mosby, Inc.
 PUBLISHER:
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AB Cardiopulmonary bypass is a potent stimulus for activation of procoagulant pathways. Heparin, the traditional antithrombotic agent, however, is often assocd. with increased perioperative blood loss because of its

English

DOCUMENT TYPE:

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multiple sites of action in the coagulation cascade and its antiplatelet
                           and profibrinolytic effects.
                         and profibrinolytic effects. Furthermore, heparin-mediated immunol. reactions (i.e., heparin-induced thrombocytopenia) may contraindicate its use. Cardiopulmonary bypass with a selective factor IXa inhibitor was tested to see whether it could effectively limit bypass circuit/intravascular space thrombosis while decreasing
                       bypass circuit/intravascular space thrombosis while decreasing extravascular bleeding, thereby providing an alternative anticoagulant strategy when heparin may not be safely administered. Active site-blocked factor IXa, a competitive inhibitor of the assembly of factor IXa into the factor X activation complex, was prepd. by modification of the enzyme's active site by the use of dansyl glutamic acid-glycine-arginine-chlormethylketone. Twenty mongrel dogs (five were given sdt. heparin/protamine: 15 were given activated site-blocked factor IXa doses ranging from 300 to 600 .mu.g/kg) underwent 1 h of hypothermic cardiopulmonary bypass, and blood loss was monitored for 3 h after the procedure. Use of activated site-blocked factor IXa as an anticoagulant in cardiopulmonary bypass limited fibrin
                        nypothermic cardiopulmonary bypass, and blood loss was monitored for 3 h after the procedure. Use of activated site-blocked factor IXa as an anticoagulant in cardiopulmonary bypass limited fibrin deposition within the extracorporeal circuit as assessed by SEM, comparable with the antithrombotic effect seen with heparin. In contrast to heparin, effective antithrombotic doses of activated site-blocked factor IXa significantly diminished blood loss in the
                    to neparin, effective antithrombotic doses of activated site-blocked factor IXa significantly diminished blood loss in the thoracic cavity and in an abdominal incisional bleeding model. These initial studies on the dog suggest that administration of activated site-blocked factor IXa may be an effective alternative anticoagulant strategy in cardiopulmonary bypass when heparin is contraindicated, affording inhibition of intravascular/extracorporeal circuit thrombosis with enhanced hemostasis in the surgical wound.

J. Thorac. Cardiovasc. Surg. (1998), 115(5), 1179-1188

CODEN: JTCSAQ: ISSN: 0022-5223

Cardiopulmonary bypass is a potent stimulus for activation of procoagulant pathways. Heparin, the traditional antithrombotic agent, however, is often assocd. with increased perioperative blood loss because of its multiple sites of action in the coagulation cascade and its antiplatelet and profibrinolytic effects. Furthermore, heparin-mediated immunol. reactions (i.e., heparin-induced thrombocytopenia) may contraindicate its use. Cardiopulmonary bypass with a selective factor IXa inhibitor was tested to see whether it could effectively limit bypass circuit/intravascular space thrombosis while decreasing extravascular bleeding, thereby providing an alternative anticoagulant
                       extravascular bleeding, thereby providing an alternative anticoagulant strategy when heparin may not be safely administered. Active
                     strategy when heparin may not be safely administered. Active site-blocked factor IXa, a competitive inhibitor of the assembly of factor IXa into the factor X activation complex, was prepd. by modification of the enzyme's active site by the use of dansyl glutamic acid-glycine-arginine-chlormethylketone. Twenty mongrel dogs (five were given std. heparin/protamine; 15 were given activated site-blocked factor IXa doses ranging from 300 to 600 .mu.g/kg) underwent 1 h of hypothermic cardiopulmonary bypass, and blood loss was monitored for 3 h after the procedure. Use of activated site-blocked factor IXa as an anticoaqulant in cardiopulmonary bypass limited fibrin
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                      hemostasis in the surgical wound.
                                                                             CAPLUS COPYRIGHT 2001 ACS
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126:31658
ACCESSION NUMBER:
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  DOCUMENT NUMBER:
TITLE:
                                                                                                     Peptide boronic acid inhibitors of trypsin-like
                                                                                                     enzymes
 INVENTOR(S):
                                                                                                     Claeson, Goran; Philipp, Manfred H. W.; Metternich,
PATENT ASSIGNEE(S):
                                                                                                     Thrombosis Research Institute, UK
SOURCE:
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                                                                                                                                                                            US 1995-459177
               R SOURCE(S): MARPAT 126:31658
Peptide boronic acids XYNHCH[(CH2)30R]BQ1Q2 (I; X = H, N-protecting group; Y = Phe-Pro; Q1Q2 = diol residue; R = Cl-4 alkyl) are inhibitors of trypsinlike enzymes (including trypsin, thrombin, factor Xa, factor IXa, factor VIIa, factor XIIa, plasmin, acrosin, complement proteases, kallikrein, urokinase, and tissue plasminogen activator), and may be administered orally or parenterally as antithrombotics. They have a rapid onset of activity and low toxicity. Thus, benzyloxycarbonyl-D-phenylalanine p-nitrophenyl ester was condensed with prollne, converted to the N-hydroxysuccinimidyl ester, coupled with the (+)-pinanediol ester of (TMS)2NCH[(CH2)3Br]B(OH)2, and reacted with quanidine-HCl and MeONa in MeOH to produce I (X = PhCH2O2C; Y = D-Phe-L-Pro; R = OMe; Q1Q2 = (+)-pinanediyl].
                                                                                                 MARPAT 126:31658
                  US 5574014 A
                                                                        19961112
                  PATENT NO.
                                                                                    KIND DATE
                                                                                                                                                                         APPLICATION NO.
                                                                                                                                                                                                                                             DATE
                                                                                                              19961112
                                                                                                                                                                         US 1994-240606
                                                                                                                                                                                                                                              19940510 <--
                 US 5856306
                                                                                                              19990105
                                                                                                                                                                         US 1995-459177
US 1998-79243
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US 6114308 A 20000905 US 1998-79243 19980514

AB Peptide boronic acids XYNHCH((CH2)3OR)BQ1Q2 (I; X = H, N-protecting group;

Y = Phe-Pro; QlQ2 = diol residue; R = Cl-4 alkyl) are inhibitors Y = Phe-Pro; QlQ2 = diol residue; R = Cl-4 alkyl) are inhibitors of trypsinlike enzymes (including trypsin, thrombin, factor Xa, factor IXa, factor VIIa, factor XIIa, plasmin, acrosin, complement proteases, kallikrein, urokinase, and tissue plasminogen activator), and may be administered orally or parenterally as antithrombotics. They have a rapid onset of activity and low toxicity. Thus, benzyloxycarbonyl-D-phenylalanine p-nitrophenyl ester was condensed with proline, converted to the N-hydroxysuccinimidyl ester, coupled with the (+)-pinanediol ester of (TMS)2NCH[(CH2)3Br]B(OH)2, and reacted with quanidine-HCl and MeONa in MeOH to produce I [X = PhCH2O2C; Y = D-Phe-L-Pro; R = OMe; QlQ2 = (+)-pinanediyl]. ANSWER 6 OF 7 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1982:223071 CAPLUS DOCUMENT NUMBER: 96:223071 Studies on oral administration of concentrated factor IX preparation Ueno, Masaharu; Horikoshi, Isamu; Takahashi, Kaoru; Sakuragawa, Nobuo Dep. Hosp. Pharm., Toyama Med. Pharm. Univ., Toyama, 930-01, Japan Yakugaku Zasshi (1902), 102(2), 202-6 CODEN: YKKZAJ; ISSN: 0031-6903 Journal

TITLE: AUTHOR (S): CORPORATE SOURCE: SOURCE: DOCUMENT TYPE: LANGUAGE: UAGE: Japanese
blood-coagulation factor IX [9001-28-9] Was stable at
4-25.degree. in mildly alk. solns. and was effectively encapsulated in
liposome prepns. contg. 5% stearylamine or 0.02 M Ca2+; oral
administration of the liposome-entrapped factor
IX shortened clotting time in dogs. The transformation of
prothrombin [9001-26-7] into thrombin [9002-04-4] was
inhibited by adding phosphatidylcholines (250 mg) or 50,000 units
aprotinin [9004-04-0]. The intestinal absorption of factor II, VII
[9001-25-6], IX, and X [9001-29-0] is described.
Studies on oral administration of concentrated factor IX
preparation Japanese preparation Yakugaku Zasshi (1982), 102(2), 202-6 CODEN: YKKZAJ; ISSN: 0031-6903 so CODEN: YKKZAJ; ISSN: 0031-6903
blood-coagulation factor IX [9001-21-9] Was stable at
4-25.degree. in mildly alk. solns. and was effectively encapsulated in
liposome prepns. contg. 5% stearylamine or 0.02 M Ca2+; oral
administration of the liposome-entrapped factor
IX shortened clotting time in dogs. The transformation of
prothrombin [9001-25-7] into thrombin [9002-04-4] was
inhibited by adding phosphatidylcholines (250 mg) or 50,000 units
aprotinin [9004-04-0]. The intestinal absorption of factor II, VII
[9001-25-6], IX, and X [9001-29-0] is described. 9001-28-9 RL: BIOL (Biological study)

(liposome-entrapped, oral administration of)

ANSWER 7 OF 7 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. JOFYRIGH J:160092 EMBASE 1987160092 Evaluati

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE: Evaluation of p-amidinophenyl esters as potential

AUTHOR . CORPORATE SOURCE:

antithrombotic agents.
Pizzo S.V.; Turner A.D.; Porter N.A.; Gonias S.L.
Department of Pathology, Duke University Medical Center,
Durham, NC 27710, United States

SOURCE: Thrombosis and Haemostasis, (1986) 56/3

(387 - 390)CODEN: THHADQ

COUNTRY: Germany

DOCUMENT TYPE: FILE SEGMENT: Journal 037

Drug Literature Index 030

SUAGE: English

Three p-amidinophenyl esters have been synthesized and characterized as irreversible inhibitors of the vitamin-K dependent proteinases; factors IXA, Xa and thrombin. In the present report we describe the in vitro and in vivo effects of these agents on standard coagulation tests in vitro and in blood from animals treated with the compounds. At a concentration of 500 .mu.M, the three esters increased the activated partial thromboplastin time (PTT) of pooled human plasma 3 to 5-fold. The prothrombin time increased 1.4 to 3.7-fold under similar conditions. The p-amidinophenyl ester of cinnamic acid (CINN) showed the most pronounced effect on both assays. This ester also is the best inhibitor of human factors IXa and Xa, while the p-amidinophenyl ester of benzoic acid (BENZ) is a slightly better alpha.-thrombin inhibitor. The effect of these esters on the thrombin clotting time correlated with in vitro kinetic measurements of .alpha.-thrombin inhibition rates.

Both BENZ and CINN increased the assay endpoint more than 6-fold. The three esters also were studied using mouse plasma. A comparable effect on the PTT was noted. Intravenous administration of 300 .mu.l of 1 mM CINN as a single bolus in mice caused a 2.3-fold increase in the PTT which remained 1.2-fold normal 2 h later. The BENZ and .alpha.-methyl-cinnamic acid (MECINN) esters were somewhat less effective as predicted from their in vitro effect on the PTT. This investigation and previous studies indicate that these compounds demonstrate low toxicity at therapeutic levels. It is concluded that the p-amidinophenyl esters may be useful in antithrombotic therapy.

Thrombosis and Haemostasis, (1986) 56/3 (387-390). English

CODEN: THHADQ
Three p-amidinophenyl esters have been synthesized and characterized as irreversible inhibitors of the vitamin-K dependent proteinases; factors IXa, Xa and thrombin. In the present report we describe the in vitro and in vivo effects of these agents on. p-amidinophenyl ester of cinnamic acid (CINN) showed the most pronounced effect on both assays. This ester also is the best inhibitor of human factors IXa and Xa, while the p-amidinophenyl ester of benzoic acid (BENZ) is a slightly better .alpha.-thrombin inhibitor. The effect of these esters on the thrombin clotting time correlated with in vitro kinetic measurements of .alpha.-thrombin inhibitor rates. Both BENZ and CINN increased the assay endpoint more than 6-fold. The three esters also were studied using mouse plasma. A comparable effect on the PTT was noted. Intravenous administration of 300 .mu.l of 1 mM CINN as a single bolus in mice caused a 2.3-fold increase in the PTT. caused a 2.3-fold increase in the PTT.

SOLOMON R?/AU

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=> s 113 and (factor IXa)
L14 89 L13 AND (FACTOR IXA)
     L14
     ≈> dup rem 114
     PROCESSING COMPLETED FOR L14
                                    37 DUP REM L14 (52 DUPLICATES REMOVED)
     => dis his
                 (FILE 'HOME' ENTERED AT 10:07:42 ON 29 MAR 2001)
              FILE 'MEDLINE, CAPLUS, EMBASE, BIOSIS' ENTERED AT 10:07:58 ON 29 MAR 2001
12853 S (FACTOR IXA) OR (FACTOR IX)
3660 S L1 (P) (MUTAT? OR INHIB?)
652552 S L2 (P) (CLOT OR THROMB?)
1125 S L2 (P) (CLOT OR THROMB?)
355 S L2 (P) (INHIBIT? (5N) (CLOT OR THROMB?))
0 S L5 AND & PPC-199609
0 S L5 AND & PPC-199609
0 S L5 AND & PPC-199609
0 S L5 AND & PPC-1999609
10 S L5 AND & PDC-1999609
7 S L5 AND & PDC-1999609
10 S L5 AND & PDC-1999609
1128 DUP REM L10 (135 DUPLICATES REMOVED)
7 S L11 (P) ADMINIST?
10574 S PINSKY D2/AU OR STERN D2/AU OR SCHMIDT A?/AU OR ROSE E?/AU OR 89 S L13 AND (FACTOR IXA)
37 DUP REM L14 (52 DUPLICATES REMOVED)
    L3
L4
L5
    L6
L7
    L8
    L10
    L12
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2 FILES SEARCHED...
           3 FILES SEARCHED.
                                  4 L15 AND PD<19960927
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    L16 ANSWER 1 OF 4 BIOSIS COPYRIGHT 2001 BIOSIS
   ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                     1991:356261 BIOSIS
                                                     BR41:40776
                                                    ACTIVE SITE-BLOCKED FACTOR IXA PREVENTS
INTRAVASCULAR CORONARY THROMBOSIS WITHOUT IMPAIRING
EXTRAVASCULAR COAGULATION.
BENEDICT C R; RYAN J; GERLACH M; WOLITZKY B; STERN
   AUTHOR(S):
                                                   DUNIV. TEXAS, HOUSTON, TEX.
JOINT MEETING OF THE ASSOCIATION OF AMERICAN PHYSICIANS,
THE AMERICAN SOCIETY FOR CLINICAL INVESTIGATION, AND THE
AMERICAN FEDERATION FOR CLINICAL RESEARCH, SEATTLE,
WASHINGTON, USA, MAY 3-6, 1991. CLIN RES, (1991) 39 (2),
   CORPORATE SOURCE:
   SOURCE:
                                                   CODEN: CLREAS. ISSN: 0009-9279.
  DOCUMENT TYPE:
FILE SEGMENT:
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   LANGUAGE:
                                                    English
  L16 ANSWER 2 OF 4
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                   BIOSIS COPYRIGHT 2001 BIOSIS 1987:58627 BIOSIS
                                                  BR32:28848
TUMOR NECROSIS FACTOR UPREGULATES FACTOR-IX-IXA BINDING
SITES AND FACTOR-IXA-VIII-MEDIATED
FACTOR-XA FORMATION ON ENDOTHELIUM.
NAWROTH P P; CORNELSON S; STERN D M
OKLA. MED. RES. FOUND., OKLAHOMA CITY, OKLA.
JOINT PROCEEDINGS OF THE 59TH SCIENTIFIC SESSIONS OF THE
AMERICAN HEART ASSOCIATION, THE 40TH ANNUAL MEETING OF THE
AMERICAN SOCIETY FOR THE STUDY OF ARTERIOSCLEROSIS (COUNCIL
ON ARTERIOSCLEROSIS), AND THE SEVENTH NATIONAL CONFERENCE
ON THROMBOSIS AND HEMOSTASIS, DALLAS, TEX., USA, NOV.
17-20, 1986. AM HEART ASSOC MONOGR, (1986) 0 (124), II-233.
CODEN: AHNOAH. ISSN: 0065-8499.
                                                   BR32:28848
   TITLE:
  AUTHOR(S):
   CORPORATE SOURCE:
   SOURCE:
  DOCUMENT TYPE:
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                                                  BR; OLD
English
  FILE SEGMENT:
  LANGUAGE:
 L16 ANSWER 3 OF 4 ACCESSION NUMBER:
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                                                 1986:194805 BIOSIS
BR30:106677
  DOCUMENT NUMBER:
 TITLE:
                                                  ACTIVATION OF COAGULATION RELEASES ENDOTHELIAL CELL
                                                  MITOGENS.
GAJDUSEK C; CARBON S; NAWROTH P; STERN D
 AUTHOR (S):
                                                 GAJDUSEK C; CARBON S; NAWROTH P; STERN D
UNIV. WASH., SEATTLE, WASH.
SYMPOSIUM ON PROTEASES IN BIOLOGICAL CONTROL AND
BIOTECHNOLOGY HELD AT THE 15TH ANNUAL UCLA (UNIVERSITY OF
CALIFORNIA-LOS ANGELES) MEETING ON MOLECULAR AND CELLULAR
BIOLOGY, LOS ANGELES, CALIF., USA, FEB. 9-15, 1986. J CELL
BIOCHEM SUPPL, (1986) 0 (10 PART A), 248.
CODEN: JCBSD7.
Conference
BR: OLD
 CORPORATE SOURCE:
  SOURCE:
 DOCUMENT TYPE:
  FILE SEGMENT:
                                                  BR; OLD
 LANGUAGE:
                                                  English
L16 ANSWER 4 OF 4
                                                BIOSIS COPYRIGHT 2001 BIOSIS
1984:77271 BIOSIS
BR26:77271
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                 THE BINDING OF FACTOR-IX AND FACTOR-IXA
TO VASCULAR ENDOTHELIAL CELLS.
STERN D M; DRILLINGS M; NOSSEL H L
DEP. MED., COLUMBIA UNIV. COLL. PHYSICIANS SURGEONS, NEW
YORK, USA.
9TH INTERNATIONAL CONGRESS ON THROMBOSIS AND HEMOSTASIS,
TITLE:
AUTHOR (S) .
CORPORATE SOURCE:
SOURCE:
                                                 JULY 4-8, 1983. THROMB HEMOSTASIS, (1983) 50 (1), 419. CODEN: THHADQ. ISSN: 0340-6245.
DOCUMENT TYPE:
                                                 Conference
FILE SEGMENT:
                                                English
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ALL L# QUERIES AND ANSWER SETS ARE DELETED AT LOGOFF LOGOFF? (Y)/N/HOLD:y

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☐ 1. Document ID: US 6200749 B1

L7: Entry 1 of 12

File: USPT

Mar 13, 2001

US-PAT-NO: 6200749

DOCUMENT-IDENTIFIER: US 6200749 B1

TITLE: Mutated forms of the ataxia-telangiectasia gene and method to screen for

a partial A-T phenotype

DATE-ISSUED: March 13, 2001

INVENTOR-INFORMATION:

NAME

CITY

STATE

ZIP CODE

COUNTRY

Shiloh; Yosef

Tel Aviv

N/A

N/A

ILX

US-CL-CURRENT: <u>435/6</u>; <u>536/23.5</u>

Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KWIC	Draw, Desc	Image

☐ 2. Document ID: US 6103244 A

L7: Entry 2 of 12

File: USPT

Aug 15, 2000

US-PAT-NO: 6103244

DOCUMENT-IDENTIFIER: US 6103244 A

TITLE: Methods for generating immune responses employing modified vaccinia of

fowlpox viruses

DATE-ISSUED: August 15, 2000

INVENTOR-INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY Dorner; Friedrich Vienna N/A N/A ATX Scheiflinger; Friedrich Orth/Donau N/A N/A ATX Falkner; Falko Gunter Mannsdorf N/A N/A ATX Pfleiderer; Michael Breitstetten N/A N/A ATX

US-CL-CURRENT: 424/199.1; 424/188.1, 424/232.1



☐ 3. Document ID: US 6093392 A

L7: Entry 3 of 12

File: USPT

PA

Jul 25, 2000

US-PAT-NO: 6093392

DOCUMENT-IDENTIFIER: US 6093392 A

TITLE: Methods and compositions for use in gene therapy for treatment of

hemophilia

DATE-ISSUED: July 25, 2000

INVENTOR-INFORMATION:

NAME

CITY

STATE ZIP CODE

COUNTRY

High; Katherine A.

Merion

N/A

N/A

Herzog; Roland W.

Glenolden

PA N/A

N/A

US-CL-CURRENT: 424/93.2; 424/93.6, 435/320.1, 435/456

Full Title Citation Front Review Classification Date Reference Claims KMC Draw Desc Image

4. Document ID: US 6046380 A

L7: Entry 4 of 12

File: USPT

Apr 4, 2000

US-PAT-NO: 6046380

DOCUMENT-IDENTIFIER: US 6046380 A

TITLE: Factor IX production in transgenic non-human mammals and factor IX DNA

sequences with modified splice sites

DATE-ISSUED: April 4, 2000

INVENTOR-INFORMATION:

NAME

CITY

STATE

ZIP CODE

COUNTRY

Clark; Anthony John

Midlothian

N/A N/A

GBX

US-CL-CURRENT: 800/14; 435/212, 435/69.6, 536/23.2, 536/23.5, 800/7

Full Title Citation Front Review Classification Date Reference Claims KMC Draw Desc Image

☐ 5. Document ID: US 6034222 A

L7: Entry 5 of 12

File: USPT

Mar 7, 2000

US-PAT-NO: 6034222

DOCUMENT-IDENTIFIER: US 6034222 A

TITLE: Method for the separation of recombinant pro-factor IX from recombinant

factor IX

DATE-ISSUED: March 7, 2000

INVENTOR-INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY Fischer; Bernhard Vienna N/A N/A ATX Mitterer; Artur Orth/Donau N/A N/A ATX Dorner; Friedrich Vienna N/A N/A ATX Eibl; Johann Vienna N/A N/A ATX

US-CL-CURRENT: <u>530</u>/<u>381</u>; <u>530</u>/<u>412</u>, <u>530</u>/<u>416</u>

Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KWIC	Draw Desc	Image
						_					-8-

☐ 6. Document ID: US 6027913 A

L7: Entry 6 of 12

File: USPT

Feb 22, 2000

US-PAT-NO: 6027913

DOCUMENT-IDENTIFIER: US 6027913 A

TITLE: Nucleic acid amplification with direct sequencing

DATE-ISSUED: February 22, 2000

INVENTOR-INFORMATION:

NAME

CITY

STATE ZIP CODE

COUNTRY

Sommer; Steven S.

Northwest Rochester

55901 MN

N/A

US-CL-CURRENT: 435/69.1; 435/91.21

Full Title Citation Front Review Classification Date Reference Claims KWC Draw Desc Image

☐ 7. Document ID: US 5891629 A

L7: Entry 7 of 12

File: USPT

Apr 6, 1999

US-PAT-NO: 5891629

DOCUMENT-IDENTIFIER: US 5891629 A

TITLE: Compositions for improving RNase cleavage of base pair mismatches in

double-stranded nucleic acids

DATE-ISSUED: April 6, 1999

INVENTOR-INFORMATION:

NAME

CITY

STATE ZIP CODE

COUNTRY

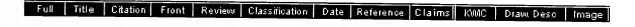
Goldrick; Marianna M.

Pflugerville

TX N/A

N/A

US-CL-CURRENT: <u>435/6</u>; <u>435/91.2</u>



■ 8. Document ID: US 5858661 A

L7: Entry 8 of 12

File: USPT

Jan 12, 1999

US-PAT-NO: 5858661

DOCUMENT-IDENTIFIER: US 5858661 A

TITLE: Ataxia-telangiectasia gene and its genomic organization

DATE-ISSUED: January 12, 1999

INVENTOR-INFORMATION:

NAME

CITY

STATE

ZIP CODE

COUNTRY

Shiloh; Yosef

Tel Aviv

N/A

N/A

ILX

US-CL-CURRENT: <u>435/6</u>; <u>536/23.5</u>

Full Title Citation Front Review Classification Date Reference Claims KMC Draw. Desc Image

9. Document ID: US 5839443 A

L7: Entry 9 of 12

File: USPT

Nov 24, 1998

US-PAT-NO: 5839443

DOCUMENT-IDENTIFIER: US 5839443 A

TITLE: Method for inhibiting thrombosis in a patient whose blood is subjected

to extracorporeal circulation

DATE-ISSUED: November 24, 1998

INVENTOR-INFORMATION:

NAME	CITY	STATE	ZIP CODE	COUNTRY
Rose; Eric	Tenafly	NJ	N/A	N/A
Stern; David	Great Neck	NY	N/A	N/A
Schmidt; Ann Marie	Franklin Lakes	NJ	N/A	N/A
Spanier; Talia	New York	NY	N/A	N/A

US-CL-CURRENT: <u>128/898</u>; <u>435/13</u>

Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KWIC	Draw Desc	Image
							1				

☐ 10. Document ID: US 5268275 A

L7: Entry 10 of 12

File: USPT

·Dec 7, 1993

US-PAT-NO: 5268275

DOCUMENT-IDENTIFIER: US 5268275 A

TITLE: Vitamin K-dependent carboxylase

DATE-ISSUED: December 7, 1993

INVENTOR-INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY Stafford; Darrel W. Carrboro NC N/A N/A Wu; Sheue-Mei . Carrboro NC N/A N/A

US-CL-CURRENT: $\frac{435}{69.1}$; $\frac{435}{232}$, $\frac{435}{252.3}$, $\frac{435}{320.1}$, $\frac{435}{352}$, $\frac{435}{354}$, $\frac{435}{358}$, $\frac{435}{366}$, $\frac{435}{69.6}$, $\frac{536}{23.2}$

Full Title Citation Front Review Classification Date Reference Claims KMC Draw. Desc Image

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Terms	Documents
(Factor adj (IXa or IX)) near mutat\$	12

Display 10 Documents, starting with Document: 11

Display Format: CIT Change Format

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Search Results - Record(s) 1 through 10 of 12 returned.

☐ 1. Document ID: US 6200749 B1

L7: Entry 1 of 12

File: USPT

Mar 13, 2001

DOCUMENT-IDENTIFIER: US 6200749 B1

TITLE: Mutated forms of the ataxia-telangiectasia gene and method to screen

for a partial A-T phenotype

DEPR:

A technical explanation for this bias towards deletions and insertions could be a greater ability of the REF method to detect such lesions versus its ability to detect base substitution. Liu and Sommer (1995) have shown, however, that the detection rate of this method in a sample of 42 point mutations in the factor IX gene ranged between 88% and 100%, depending on the electrophoresis conditions. The 7 base substitutions detected directly by the REF method in the present study (Table 2), indicate that such sequence alterations are detected in our hands as well.

	The real Party lies	-									
Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KONAC	Draini Desc	Image
			_							0.000	1111555

☐ 2. Document ID: US 6103244 A

L7: Entry 2 of 12

File: USPT

Aug 15, 2000

DOCUMENT-IDENTIFIER: US 6103244 A

TITLE: Methods for generating immune responses employing modified vaccinia of

fowlpox viruses

DEPR:

Human clotting factor IX is a 56 kDa glycoprotein involved in the regulation of blood coagulation. This clotting factor undergoes complex post-translational modifications: vitamin K dependent carboxylation of the first 12 glutamic residues, glycosylation, 3-hydroxylation of an aspartic acid and amino terminal protein processing. Davie, E. W., "The Blood Coagulation Factors: Their cDNAs, Genes and Expression", HEMOSTATIS AND THROMBOSIS, Colman et al., eds., J. B. Lippincott Co. (1987). Hemophilia B, an X chromosome-linked bleeding disorder, is caused by mutation of factor IX. Patients with hemophilia are currently treated by substitution with plasma-derived factor IX.

								· · · · · · · · · · · · · · · · · · ·			
Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KOME:	Drami Desc	Image
										B1000 B 000	1111282

☐ 3. Document ID: US 6093392 A

L7: Entry 3 of 12

File: USPT

Jul 25, 2000

DOCUMENT-IDENTIFIER: US 6093392 A

TITLE: Methods and compositions for use in gene therapy for treatment of

hemophilia

BSPR:

In another aspect, the isolated DNA encoding Factor IX comprises a mutation which renders Factor IX encoded thereby incapable of binding to collagen IV.

BSPR:

In yet another aspect, the isolated DNA encoding Factor IX comprises a mutation which renders Factor IX encoded thereby incapable of binding to collagen IV.

CLPR:

7. The method of claim 1, wherein said nucleic acid encoding Factor IX comprises a mutation which reduces binding of Factor IX encoded thereby to collagen IV as compared to a Factor IX lacking the mutation, wherein the mutation replaces a lysine residue with an alanine residue in the fifth amino acid position from the beginning of mature Factor IX.



☐ 4. Document ID: US 6046380 A

L7: Entry 4 of 12

File: USPT

Apr 4, 2000

DOCUMENT-IDENTIFIER: US 6046380 A

TITLE: Factor IX production in transgenic non-human mammals and factor IX DNA sequences with modified splice sites

ORPL:

Chen, S.-H. et al., "Splice junction mutations in factor IX gene resulting in severe hemophilia B, " Nucl. Acids Res. 19(5):1172 (1992).

Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KWIC	Draw, Desc	image

5. Document ID: US 6034222 A

L7: Entry 5 of 12

File: USPT

Mar 7, 2000

DOCUMENT-IDENTIFIER: US 6034222 A

TITLE: Method for the separation of recombinant pro-factor IX from recombinant

factor IX

BSPR:

Up to now, an improvement in the recovery of recombinant, physiologically active Factor IX could only be achieved through genetic manipulation of the pro-sequence. It has thus been attempted to couple the pro-sequence of Factor VII to the DNA sequence of Factor IX in order to obtain a more effective cleavage of the pro-sequence (K. Berkner et al., Current Advances in Vitamin K Research, Elsevier Science Publishing Co., Inc. (1988) 199-207). P. Meulien et al., Prot. Engineer. 3 (1990) 629-633) examined the influence of mutations in the region of the pro-peptide cleavage site of Factor IX. They determined that the yield of active Factor IX can be distinctly increased by introduction of a point mutation in position +1 (alanine versus tyrosine); in comparison with wild-type Factor IX, which demonstrates a specific activity of 45-55% after purification over a DEAE-Sepherodex.RTM. column and stepwise elution with 0.3 M NaCl in the physiological pH range, a specific activity of 85 to 100% was found for the mutated Factor IX.

Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KWIC	Draw Desc	Image

☐ 6. Document ID: US 6027913 A

L7: Entry 6 of 12

File: USPT

Feb 22, 2000

DOCUMENT-IDENTIFIER: US 6027913 A

TITLE: Nucleic acid amplification with direct sequencing

DEPR:

Direct sequencing also makes it feasible to delineate point mutations in multiple individuals. For an X-linked lethal disease, direct sequencing can provide a "snapshot" of recent mutations in the population because the mutations that arise are extinguished within a few generations [Haldane, J. B. S., Genet, 31:317-326 (1935)]. Analysis of such data should reveal whether any hotspots of mutation exist. Previously protein and nucleic acid sequence of hundreds of variant .alpha.- and .beta.- globin alleles did not reveal any dramatic hotspots in these autosomal genes [Vogel, F. and A. G. Motulsky (eds) In: Human Genetics. Edition 2, Springer-Verlag, Berlin, pp. 433-511, (1986)). Notably, transitions of CpG were not markedly elevated. More recently the delineation of mutations in other genes has indicated that transitions at CpG occur with great frequency [Youssoufian, H., H. H. Kazazian, Jr., D. B. Phillips, S. Aronis, G. Tsifitis, V. A. Brown, S. E. Antonarkis, Nature, 324:380-382 (1986); Youssoufian, H., S. E. Antonarakis, W. Bell, A. M. Griffin, H. H. Kazazian, Jr., Am. J. Hum. Genet., 42:718-725 (1988); Vulliamy T. J., M. D. Urso, G. Battistuzzi, M. Estrada, N. S. Foulkes, G. Martini, V. Calabro, V. Poggi, R. Giordana, M. Town, L. Luzzato, M. G. Persico, Proc. Natl. Acad. Sci. USA, 85:5171-5175 (1988); Cooper D. N. and H. Youssoufian, Hum. Genet., 78:151-155 (1988)]. Eight regions of likely functional significance in 21 hemophiliacs from different families have been sequenced. The results of this large sample of germline mutations from a single gene show that CpG is a hotspot of $\underline{\text{mutation in the factor IX}}$ gene and that the rate of enhancement is about 77-fold. This enhancement is not restricted to a particular subset of CpGs with constant bases in the immediately flanking sequence.



☐ 7. Document ID: US 5891629 A

L7: Entry 7 of 12

File: USPT

Apr 6, 1999

DOCUMENT-IDENTIFIER: US 5891629 A

 ${\tt TITLE:}$ Compositions for improving RNase cleavage of base pair mismatches in double-stranded nucleic acids

BSPR:

Of the three $\frac{\text{mutations}}{\text{A}}$ in the Factor IX model system that were not detected by either RNase A alone or the RNase A/RNase I combination used in the initial studies, all are detected using the new RNase digestion conditions disclosed in the present invention. Mutations in the p53 tumor suppressor gene that are not detected by RNase A are also detected by RNase I using the new conditions (FIG. 6A, FIG. 6B, and FIG. 6C). Moreover, when the entire panel of 60 mismatches in the model system (2 complementary mismatches are generated from each of the 30 point mutations) is compared using the new RNase digestion components and the components used in the method described in U.S. patent application Ser. No. 08/371,531, it is clear that the new conditions show a dramatic improvement in the general ability to specifically cleave a wide variety of mismatches.

DRPR:

FIGS. 7A-7B: Cleavage of mismatches in a large panel of homozygous and heterozygous samples with $\underline{Factor\ IX\ mutations}$. FIG. 7 is composed of two panels: FIG. 7A and FIG. 7B.

DEPR:

This example details the cleavage and detection of mismatches with digestion buffers of the present invention in large panel homozygous and heterozygous samples with Factor IX mutations. Double-stranded RNA targets containing mismatches due to point mutations in exon 8 of the Factor IX gene were prepared from genomic DNA isolated from Hemophilia B patients and heterozygous carriers, as described in Example 2.

DEPC:

Cleavage of Mismatches in Large Panel of Homozygous and Heterozygous Samples with Factor IX Mutations

Full Title Citation Front Review Classification Date Reference Claims KWC Draw Desc Image

■ 8. Document ID: US 5858661 A

L7: Entry 8 of 12

File: USPT

Jan 12, 1999

DOCUMENT-IDENTIFIER: US 5858661 A

TITLE: Ataxia-telangiectasia gene and its genomic organization

A technical explanation for this bias towards deletions and insertions could be a greater ability of the REF method to detect such lesions versus its ability to detect base substitution. Liu and Sommer (1995) have shown, however, that the detection rate of this method in a sample of 42 point mutations in the factor IX gene ranged between 88% and 100%, depending on the electrophoresis conditions. The 7 base substitutions detected directly by the REF method in the present study (Table 2), indicate that such sequence alterations are detected in our hands as well.



☐ 9. Document ID: US 5839443 A

L7: Entry 9 of 12

File: USPT

Nov 24, 1998

DOCUMENT-IDENTIFIER: US 5839443 A

TITLE: Method for inhibiting thrombosis in a patient whose blood is subjected to extracorporeal circulation

DEPU:

Wacey, A. I., Krawczak, M., Kakkar, V. V. and Cooper, D. N. (1994) Determinants of the factor IX mutational spectrum in haemophilia B: an analysis of missense mutations using a multi-domain molecular model of the activated protein. Hum. Genet. 94:594-608.



☐ 10. Document ID: US 5268275 A

L7: Entry 10 of 12

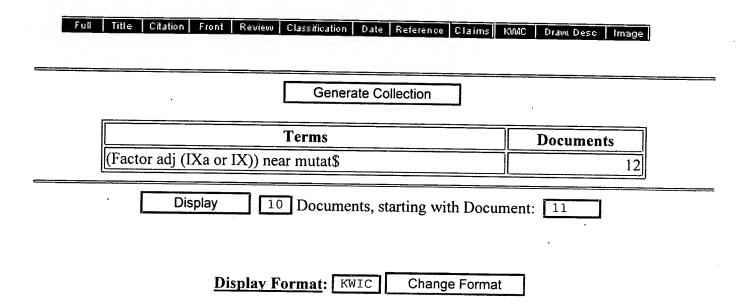
File: USPT

Dec 7, 1993

DOCUMENT-IDENTIFIER: US 5268275 A TITLE: Vitamin K-dependent carboxylase

DEPR:

Preparation of Affinity Column. Peptide FIXQ/S SEQ ID NO:1) (residues -18 to 41 of factor IX with mutations Arg to Glu at residue -4 and Arg to Ser at residue -1) was chosen for the affinity ligand because its affinity for the carboxylase is not changed and because it has fewer trypsin cleavage sites than our other peptides and is therefore less likely to be degraded by proteases in the crude extracts used for purification. Peptide FIXQ/S was prepared according to S.-M. Wu et al. supra. One hundred mg of FIXQ/S was coupled to 25 ml of Affi-Gel 10 (Bio-Rad Inc.) according to the manufacturer. The reaction was done at pH 4.8, which is one unit below the theoretical pI of FIXQ/S. The final concentration of the covalently bound FIXQ/S on Affi-Gel 10 was measured as 442 .mu.M and the coupled ligand is referred to as Affi-FIXQ/S.



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☐ 11. Document ID: US 4994371 A

L7: Entry 11 of 12

File: USPT

Feb 19, 1991

US-PAT-NO: 4994371

DOCUMENT-IDENTIFIER: US 4994371 A

TITLE: DNA preparation of Christmas factor and use of DNA sequences

DATE-ISSUED: February 19, 1991

INVENTOR-INFORMATION:

NAME

CITY

STATE

ZIP CODE

COUNTRY

Davie; Earl W.

Bellevue

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98004

N/A

Kurachi; Kotoku

Seattle

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98125

N/A

US-CL-CURRENT: $\frac{435}{6}$; $\frac{435}{243}$, $\frac{435}{243}$, $\frac{435}{320.1}$, $\frac{435}{91.41}$, $\frac{435}{91.51}$, $\frac{436}{501}$, $\frac{436}{504}$, $\frac{536}{23.5}$, $\frac{536}{24.31}$, $\frac{536}{25.32}$

Full	Title	Citation	Front	Review	Classification	Date	Reference	Claims	KWIC	Draw Desc Image
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☐ 12. Document ID: CA 2002540 C, EP 373012 A, FR 2638643 A, CA 2002540 A, JP 02265487 A, EP 373012 B1, DE 68920980 E, US 5521070 A, JP 2936201 B2

L7: Entry 12 of 12

File: DWPI

Apr 4, 2000

DERWENT-ACC-NO: 1990-180758

DERWENT-WEEK: 200035

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TITLE: DNA coding for human factor IX - with mutation in pro coding sequence

INVENTOR: MEULIEN, P

PRIORITY-DATA: 1988FR-0014635 (November 9, 1988)

PATENT-FAMILY:

PUB-NO	PUB-DATE	LANGUAGE	PAGES	MAIN-IPC
CA 2002540 C	April 4, 2000	F	000	C12N015/57
EP 373012 A	June 13, 1990	N/A	000	N/A
FR 2638643 A	May 11, 1990	N/A	000	N/A
CA 2002540 A	May 9, 1990	N/A	000	N/A
JP 02265487 A	October 30, 1990	N/A	000	N/A
EP 373012 B1	February 1, 1995	F	011	C12N015/57
DE 68920980 E	March 16, 1995	N/A	000	C12N015/57
US 5521070 A	May 28, 1996	N/A	007	C12N015/00
JP 2936201 B2	August 23, 1999	N/A	012	C12N015/09

INT-CL (IPC): A61K 31/00; A61K 37/54; A61K 38/43; A61K 38/46; C07H 21/04; C07K 14/00; C07K 15/06; C12N 5/10; C12N 9/64; C12N 15/00; C12N 15/09; C12N 15/57; C12N 15/86; C12P 9/00; C12P 21/00; C12P 21/02; C12P 21/06; C12R 1/91

Full Title	e Citation	Front	Review	Classification	Date	Reference	Claims	KWIC	Drawu Desc	Image
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	,			Terms					Document	S

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Search Results - Record(s) 11 through 12 of 12 returned.

☐ 11. Document ID: US 4994371 A

L7: Entry 11 of 12

File: USPT

Feb 19, 1991

DOCUMENT-IDENTIFIER: US 4994371 A

TITLE: DNA preparation of Christmas factor and use of DNA sequences

DEPR:

In accordance with the subject invention, DNA sequences are provided for hybridization with pro-factor IX, factor IX, factor IX.sub.a, and activation peptide, and for DNA and RNA fragments which can be used in the detection of mutations or other genetic deficiencies concerned with factor IX. The sequences can be used in diagnosing blood clotting deficiencies, such as hemophilia, particularly hemophilia B. By lysing cells as described above and screening the DNA with fragments according to the subject invention, mutations in the factor IX gene may be determined.



12. Document ID: CA 2002540 C, EP 373012 A, FR 2638643 A, CA 2002540 A, JP 02265487 A, EP 373012 B1, DE 68920980 E, US 5521070 A, JP 2936201 B2

L7: Entry 12 of 12

File: DWPI

Apr 4, 2000

DERWENT-ACC-NO: 1990-180758

DERWENT-WEEK: 200035

COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: DNA coding for human factor IX - with mutation in pro coding sequence

Full Title Citation Front Review Classification Date Reference Claims KWIC

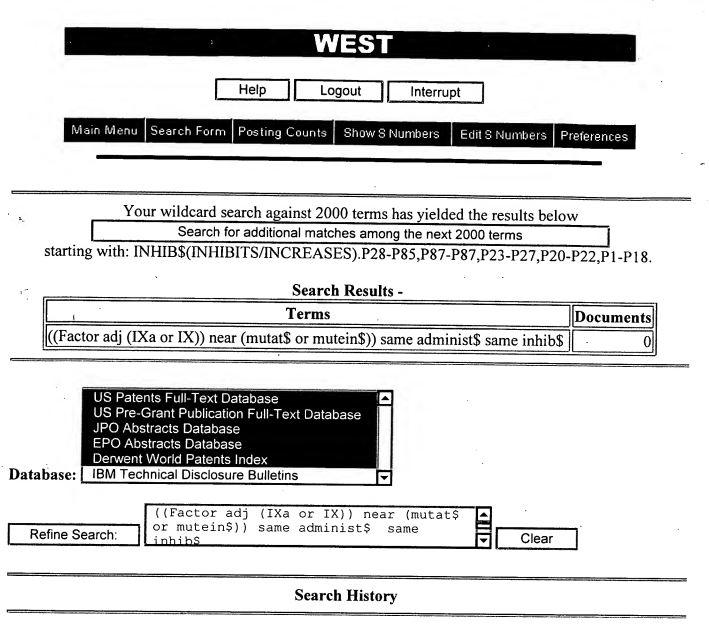
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USPT,PGPB,JPAB,EPAB,DWPI	((Factor adj (IXa or IX)) near (mutat\$)) same administ\$ same inhib\$	0	<u>L5</u>
USPT,PGPB,JPAB,EPAB,DWPI	((Factor adj (IXa or IX)) near (mutat\$)) same administ\$ same (clot or thromb\$)	0	<u>L4</u>
USPT,PGPB,JPAB,EPAB,DWPI	((Factor adj (IXa or IX)) near (mutat\$)) near administ\$ same (clot or thromb\$)	0	<u>L3</u>
USPT,PGPB,JPAB,EPAB,DWPI	(Factor adj (IXa or IX)) near (mutat\$) near administ\$ near (clot or thromb\$)	0	<u>L2</u>
USPT,PGPB,JPAB,EPAB,DWPI	(Factor adj (IXa or IX)) near (mutat?) near administ? near clot	0	<u>L1</u>

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L10 ANSWER 8 OF 15 MEDLINE
     . . Russell's Viper Venom time and by the Textarin/Ecarin ratio.
AΒ
     APC-response was studied by a clotting (aPTT-based) and by an amidolytic
     factor IXa-X-based) assay. A reduced response to APC
     (APC-resistance) was found in 49% of 65 PLa-positive and in 13% of 90
     PLa-negative. . . common in the samples with LA, as compared to
CLa+PSa
     positive (58% vs. 30%, not significant). The presence of the
     mutation causing Arg506-Gln substitution in coagulation factor V
     was investigated in 84 samples. The occurrence of the mutation
     in APC-resistant patients with CLa+PSa or with LA in one of the two
assavs
     was similar to those without PLa (84% and 100%, respectively). In the
     absence of APC resistance, the occurrence of the mutation was
     similar in the samples with and without PLa (14% vs. 11%). Samples with
     LA, determined by both tests used, comprised a special group where the
     frequency of the mutation in the APC resistant samples was
     significantly reduced (p < 0.01). In the latter samples, the pathogenic
    mechanism of APC.
V Deficiency: BL, blood
     *Factor V Deficiency: GE, genetics
      Factor V Deficiency: IM, immunology
      IgG: IM, immunology
      IgM: IM, immunology
     Lupus Coagulation Inhibitor: AN, analysis
     Middle Age
     Molecular Sequence Data
     *Partial Thromboplastin Time
      Phosphatidylserines: IM, immunology *
     *Phospholipids: IM, immunology
    0 (Antibodies, Anticardiolipin); 0 (Antibodies, Antiphospholipid); 0
     (IgG); 0 (IgM); 0 (Lupus Coagulation Inhibitor); 0
     (Phosphatidylserines); 0 (Phospholipids); 0 (Protein C)
L10 ANSWER 9 OF 15 MEDLINE
    To elucidate the role of the P1' residue of the serpin, antithrombin
AB
(AT),
    in proteinase inhibition, the source of the functional defect in
    a natural Ser-394-->Leu variant, AT-Denver, was investigated. AT-Denver
    inhibited thrombin, Factor IXa, plasmin, and
    Factor Xa with second order rate constants that were 430-, 120-, 40-, and
    7-fold slower, respectively, than those of native AT, consistent with an
    altered specificity of the variant inhibitor for its target
    proteinases. AT-Denver inhibited thrombin and Factor Xa with
    nearly equimolar stoichiometries and formed SDS-stable complexes with
    these proteinases, indicating that the diminished inhibitor
    activity was not due to an enhanced turnover of the inhibitor as
    a substrate. Binding and kinetic studies showed that heparin binding to
    AT-Denver as well as heparin accelerations of AT-Denver-proteinase
    reactions were normal, consistent with the P1' mutation not
    affecting the heparin activation mechanism. Resolution of the two-step
    reaction of AT-Denver with thrombin revealed that the majority of.
    was localized in the second reaction step and resulted from a 190-fold
    decreased rate constant for conversion of a noncovalent proteinase-
    inhibitor encounter complex to a stable, covalent complex. Little
    or no effects of the mutation on the binding constant for
```

encounter complex formation or on the rate constant for stable complex dissociation were evident. These. . . . a role for the Pl' residue of antithrombin in transition-state stabilization of a substrate-like attack of the proteinase on the inhibitor-reactive bond following the formation of a proteinase-inhibitor encounter complex but prior to the conformational change leading to the trapping of proteinase in a stable, covalent complex. Such. . .

L10 ANSWER 10 OF 15 MEDLINE

The purpose of this study is to determine which residues of the AΒ factor IXa heavy chain are important for interaction with the cofactor of factor IXa, factor VIIIa. Because the monoclonal antibody (MoAb) FXC008 inhibits interaction between factors IXa and VIIIa, and because it also reacts with residues 181-310 of the factor IXa heavy chain, we used the computer-modelled structure of the factor IXa heavy chain to select charged surface residues likely to interact with FXC008 and/or factor VIIIa. We made mutations in the region of residues 181-310 of the heavy chain of factor IX, and replaced these amino acids individually with those located at the same position in factor X. The mutated factor IX retained complete clotting activity and thus interacted normally with factor VIIIa. Five mutant proteins (factor IXK214F, factor IXK228R,. . . IXD276K nor factor IXR248H bound to FXC008. Factor IXR252V had reduced affinity to FXC008. Our results suggest the following: (1) factor IXa residues 214, 228, 240, 247, 248, 252, 260, and 276 are not involved in specific interaction with factor VIIIa; and.

L10 ANSWER 11 OF 15 MEDLINE

AB Inherited resistance to activated protein C (APC) is a recently identified

major cause of thrombosis. It is associated with a **mutation** in the factor V gene affecting one of the cleavage sites for APC. APC resistance was recently found to be. . . in a purified system. The APC-mediated degradation of factor VIIIa was monitored by a factor X activation reaction using purified **factor IXa**, phospholipid, and calcium. In the presence of both factor V and protein

APC was found to inhibit factor VIIIa activity efficiently. APC alone or together with factor V was ineffective, whereas APC in combination with protein S. . . in the reaction. Two monoclonal antibodies, one against protein S and the other directed toward factor V, were found to inhibit the APC cofactor activity of the factor V-protein S mixture. Factor Va did not express APC cofactor activity, and addition of excess factor Va over factor V did not inhibit the APC cofactor function of a factor V-protein S mixture. In conclusion, the results suggest that factor V and protein. . .

L10 ANSWER 12 OF 15 MEDLINE

AB Factor IX is a multidomain protein and is the proenzyme of a serine protease, factor IXa, essential for hemostasis. In this report, we describe the molecular basis of hemophilia B (deficiency of factor IX activity) in. . . rearrangements of the factor IX gene. By

enzymatic amplification and sequencing of all exons and promoter regions, the following causative mutation in the protease domain of factor IX was identified in each patient: IXSchmallenberg: nucleotide 31,215G---T, Ser365Ile; IXVarel: nucleotide 31,214A---G, Ser365Gly;... Arg248Gln; and IXMonschau: nucleotide 30,855A---T, Glu245Val. In IXVarel, nucleotide 31,213T was also replaced by C, which results in a silent mutation (GAT---GAC) at Asp-364. Thus, this patient has a double base-pair substitution of TA to CG at nucleotides 31,213 and 31,214. . . 40% to 100% except for IXDreihacken (Arg248Gln), in which case it was approximately 4% of normal. The Ser365Gly and Ser365Ile mutants are nonfunctional because of lack of the active site serine residue. Mutant Asp364His is inactive because it cannot form the.

observed in other homologous serine proteases, this hydrogen bond is essential for maintaining the correct active site conformation in normal factor IXa (IXaN). Purified Arg248Gln had approximately 41% and Glu245Val had approximately 17% of the activity of normal factor IX (IXN) in. . . mutant did and the Arg248Gln mutant did not bind to an anti-IXN monoclonal antibody that has been shown previously to inhibit the interaction of factor VIIIa with factor IXaN. We have recently shown that a high-affinity calcium binding site exists in. L10 ANSWER 13 OF 15 MEDLINE AB . . . IX. Also, after treatment with factor XIa, none of the Bm variants reacted with antithrombin III (in contrast to normal factor IXa). Purified factor IX Deventer (one of the variants with a replacement of Arg181), either with or without pretreatment with factor XIa, was found to be a more effective competitive inhibitor of the factor VIIa-tissue factor-induced factor X activation than similarly treated normal factor IX. In addition, this inhibitory effect was much more pronounced when bovine tissue factor was used instead of human tissue factor. We propose that the. site serine that allows efficient substrate binding and catalysis, but that the same conformational change is instrumental in effectively dissociating factor IXa from the activating factor VIIa-tissue factor complex. Amino acid replacements that disrupt this conformational transition directly (e.g. Pro368----Thr near the catalytic center) or indirectly (mutations at the Arg180-Val activation site) therefore lead to a combination of 1) the loss of coaqulant and 2) an inhibitory effect in the ox brain prothrombin time assay. *Factor IX: GE, genetics Factor IX: IM, immunology Factor IX: ME, metabolism Factor IX: PD, pharmacology Factor VIIa: ME, metabolism Factor X: AI, antagonists & inhibitors Factor XIa: ME, metabolism *Hemophilia A: BL, blood Molecular Sequence Data Molecular Weight *Mutation L10 ANSWER 14 OF 15 MEDLINE Factor IX is the zymogen of the serine protease factor IXa involved in blood coagulation. In addition to a catalytic domain homologous to the chymotrypsin family, it has Ca2+, phospholipid, . . and beta-OH aspartic acid content, and in its binding to an anti-IXN monoclonal antibody which has been shown previously to inhibit the interaction of factor VIIIa with factor IXaN. Further, IXER is cleaved to yield a factor IXa-like molecule by factor XIa/Ca2+ at a rate similar to that observed for IXN. However, in contrast to IXaN, IXaER does not bind to antithrombin-III (specific inhibitor of IXaN) and does not catalyze the activation of factor X (substrate) to factor Xa. To identify the mutation in IXER, all eight exons of IXN and IXER gene were amplified by the polymerase chain reaction technique and cloned. A single point mutation (G----T) which results in the replacement of Val for Gly363 in the catalytic domain of IXER was identified. Gly363 in factor IXa corresponds to the universally conserved Gly193 in the active

site sequence of the chymotrypsin serine protease family. X-ray crystallographic data. . . Our computer structural data support a

concept that the Gly363----Val change prevents the development of the active site conformation in **factor IXa** such that the substrate binding site and the oxyanion hole are not formed in the **mutated** enzyme.

- L10 ANSWER 15 OF 15 MEDLINE
- TI Replacement of isoleucine-397 by threonine in the clotting proteinase factor IXa (Los Angeles and Long Beach variants) affects macromolecular catalysis but not L-tosylarginine methyl ester hydrolysis. Lack of correlation between the ox brain prothrombin time and the mutation site in the variant proteins.
- AB . . . two non-functional Factor IX variants, namely Los Angeles (IXLA) and Long Beach (IXLB). Both variants could be cleaved to yield Factor IXa-like molecules, but were defective in catalysing the cleavage of Factor X (macromolecular substrate) and in binding to antithrombin III (macromolecular inhibitor). In the present study we have identified the mutation of IXLA by amplifying the exons (including flanking regions) as well as the 5' end of
- the gene by polymerase-chain-reaction. . . substitution (T----C) in exon VIII of IXLA, with a predicted replacement of Ile-397 to Thr in the mature protein. This mutation is the same as found recently for IXLB. The observation that IXLB and IXLA have the same mutation is an unexpected finding, since, on the basis of their ox brain prothrombin time (PT, a test that measures the ability of the variant Factor IX molecules to inhibit the activation of Factor X by Factor VIIa-tissue factor complex), these variants have been classified into two different groups and. . . thought to be genetically different.

Our observation thus suggests that the ox brain PT does not reflect the locus of mutation in the coding region of the variant molecules. However, our analysis suggests that the ox brain PT is related to Factor IX antigen concentration in the patient's plasma. Importantly, although the mutation in IXLA or IXLB protein is in the catalytic domain, purified IXaLA and IXaLB hydrolyse L-tosylarginine methyl ester at rates.

. . data on Factor IXBm Lake Elsinore (Ala-390----Val mutant), strengthen a conclusion that the peptide region containing residues 390-397 of normal Factor IXa plays an essential role in macromolecular substrate catalysis and inhibitor binding. However, the two mutations noted thus far in this region do not distort S1 binding site in the Factor IXa enzyme.